PATENT SPECIFICATION

(11) **1 436 502**

36 502

5

15

20

25

30

35

(21) Application No. 43098/74 (22) Filed 4 Oct. 1974

(31) Convention Application No. 419 319

(32) Filed 4 Oct. 1973 in

(33) Spain (ES)

(44) Complete Specification published 19 May 1976

(51) INT CL² C07D 211/80; A61K 27/00; C07D 231/06, 277/08, 277/20

(52) Index at acceptance

C2C 1382 1384 1401 1530 215 220 226 22Y 250 251 252 256 25Y 281 29X 29Y 30Y 342 34Y 351 355 363 36Y 603 623 625 62X 672 699 790 79Y KR KS

(72) Inventors ROBERT G. W. SPICKETT, ARMANDO VEGA NOVEROLA and JOSE PRIETO SOTO



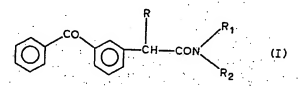
20

(54) AMIDE DERIVATIVES OF 3-BENZOYL-PHENYLALKANOIC ACIDS

(71) We, ANTONIO GALLARDO S.A., of Cardoner 68—74, Barcelona 12, Spain, a body corporate organised under the laws of Spain, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

This invention relates to new amide derivatives of 3-benzoylphenylalkanoic acids which have anti-inflammatory, analgesic and antipyretic activity and are of value for the treatment inter alia of painful inflammatory conditions such as rheumatoid arthritis, osteoarthritis and various non-specific types of inflammatory disease affecting fibromuscular tissue. The invention also relates to pharmaceutical compositions comprising the new derivatives.

According to one aspect of our invention, we provide a compound corresponding to the general formula (I):



where R represents a hydrogen atom, lower alkyl (C₁—C₂) radical or cycloalkyl radical; R₁ represents a hydrogen atom or lower (C₁—C₂) alkyl radical; and R₂ represents a heterocyclic group having one or more heteroatoms, or R₁ and R₂ together with the adjoining nitrogen atom form 3-0x0-4,5-benzo-1,2-thiazolinyl-1,1-dioxide, or a pharmaceutically acceptable salt thereof.

The radical R in formula (I) is preferably a hydrogen atom or a methyl group. R, is preferably a hydrogen atom. R₂ is preferably 2-thiazolinyl, 4-methylpyridyl, 3-hydroxypyridyl, pyridyl, 1,5-dimethyl-2-phenyl-pyrazolonyl, or thiazolyl.

According to another aspect of our invention, we provide a pharmaceutical composition comprising a compound of formula (I) as defined above, together with a non-toxic pharmaceutically acceptable carrier or diluent therefor.

The carrier or diluent may be solid or liquid. Preferred examples are lactose, corn starch, colloidal silicon dioxide, microcrystalline cellulose, carboxymethyl starch, hydroxypropyl cellulose, magnesium stearate and adeps solidus.

According to a further aspect of our invention, we provide a process for preparing a compound of formula (I) as defined above, which comprises hydrolysing a 3-benzoylphenyl α -substituted acetonitrile to form the corresponding alkanoic acid, converting the acid to an active derivative, and reacting the active derivative with an amine to form the desired amide derivative of formula (I).

The compounds may be prepared from the corresponding 3-benzoylphenyl α -substituted acetonitrile by hydrolysis in aqueous mineral acids, such as sulphuric, hydrochloric or phosphoric acid or organic acids such as formic, acetic, halogen substituted acetic acids or propionic acid at temperatures in the range of from 70° to 100°C.,

5

10

5

10

when the corresponding 3-benzoylphenyl alkanoic acids are obtained. These may be converted into the acid chlorides in solvents such as benzene, toluene, chloroform or xylene with chlorinating agents such as thionyl chloride, phosphorus pentachloride or oxalylchloride at temperatures in the range of from 80° to 120°C. The acid chlorides may then be reacted with an amine of the general formula (II).

IN R₂

in which R₁ and R₂ have the same meaning as indicated above, in solvents such as benzene, toluene, acetone or dioxane and in the presence of a strong base such as sodium hydroxide, potassium hydroxide, triethylamine or pyridine. The reaction is controlled and maintained at room temperature initially and finally completed at 70°—90°C.

In the screening tests used to detect antiinflammatory, analgesic and antipyretic activity, some of the compounds were highly active and were shown to be intermediate in activity between the known antiinflammatory agents, phenylbutazone and indomethacin. The activity of some of the compounds is shown below:

Struc	ture I				
R R ₁	R ₂		*carrag. oedema	*analgesic activity	*antipyret. activity
	·	CH ₃	•		
н н			4	4	4
СН, Н	{s	•	5	5	5
* .	N				
н н	s N		4	4	4
	,c	CH ₃			95 4
СН, Н			4	4	4
Phenylbutazone			3	1	1
Indomethacine			5	5	5

^{*} Activity is expressed as approximate ED_{50} values (mg/kg. per os) as follows >250 = 0; 126-250 = 1; 63-125 =2; 31-62 =3; 15-30 = 4, <15 =0.

	1,436,502	3
	Those compounds having a sufficiently basic heteratom may be used in the form of salts with organic or inorganic acide	
	The organic of more and action	
	For the preparation of pharmaceutical compositions the active compounds may be diluted with pharmaceutically acceptable ingredients to form the compositions of this invention, the type of excipients used dispersions to form the compositions of	
5	The true of all the true of true of the true of true of the true of the true of true o	
		5
	a series of prepared using exchinents known in the art for this doctor farms	*
10	The pharmaceutical formulations may contain from 25 to 300 mg. and the daily dose of the active component may vary between 20 mg. and 1000 mg. per day.	
	THE POLICE LABINDLES HINSITATE THE INVENTION OF COME DISCOURT AND A THE PARTY OF THE PROPERTY	10
	concern the production of intermediate compounds.	
	Example 1.	
15	3-benzovlphenyl acetic acid (Intermediate Company)	
13	11 miature of 3-benzovibnenvi acetonitrile (50 g \ water (50 -1)	15
	(50 ml) and concentrated sulphuric acid (50 ml) was refluxed with agitation for 2	12
	After cooling the mixture was noured into motor and extended with	
20		
	The parties octors cyapolanily life solvent the pasidizat colid was an all a it	20
	with benzene and dried to give a yield of 32 g. m.p. 112-4°C.	~~
	Example 2.	
	α-(3-benzovlphenyl) propionyl chloride (Intermediate Compound)	
25	" " " " " " " " " " " " " " " " " " "	•
	treated with thionyl chloride (2.5 ml.) and the solution was refluxed for 6 hours. The solvent was removed in vacuo and the residue was redissolved in benzene and evaporated to dryness. This operation was redissolved in benzene and evaporated to dryness.	25
	The second and application was remained contend towar to second at	
. **	thionyl chloride. In this way, a light coloured oil was obtained (5.3 g), which was used for the following reactions.	
	101 To lond and leactions,	
30 .	Example 3.	20
	2-[α-(3-benzoylphenyl) propionamide]-4-methyl pyridine	30
	2-amino-4-methyl pyridine (8 g. 0.04 moles) and triethylamine (4 g. 0.04 moles) were dissolved in dioxane (50 ml). To the space of 1/2 hours of 1/2	•
	of the space of 1/2 Hour, q=(3-penzovinnenvi) propional chloride (11 a	
35	ord mores) dissolved in dioxane (20 ml). When the addition was completed the min	35
	the was meated at ou C. for 2 hours. The mixture was notired into ice waser and an	, ,
	tracted several times with methylene chloride. The extracts were washed successively with water, bicarbonate and water until the washings were neutral. The extract was dried over sodium culphase and the	· · · ·
40	The	
40	hydrochloride was prepared by treating an ethanolic solution of the product with HCl, m.p. 180—182°C.	40
	Evenule 4	
٠.	By the procedure described in Example 3, the following amides more	
15	and appropriate send cittorines sind smines:	
	3-(3'-benzoylphenyl acetamido)-2-thiazoline — m.p. 161—62°C. 2-(3'-benzoylphenyl acetamido)-thiazole — m.p. 168—70°C.	45
٠. ٠.	2-(3-benzoyipnenyi acetamido)-4-methyl puridine — m n 86—88°C	•
٠.	2-(3 -belizoyiphenyi acetamido)-3-hydroxynyridine — m n 144coc	
0	2-(3'-benzoylphenyl acetamido)-3-oxo-4,5-benzo-1-2-thiazoline-1,1-dioxide — m.p, 156—57°C.	
	3-(3'-benzoylphenyl acetamido)-pyridine — m.p. 108—109C	50
· .	7-(3 -benzoyipnenyi acetamido)-1.5-dimethyl-2-phenyl pyrazoline - mp. 165 700	•
	TO THE TENED THE PROPERTY OF T	
5 . ,	4-[a(3-henzoviphenyi)propionamide]-iniazole — m.p. 130—1°C.	EE
	chloride m.p. 140—2°C. hydro-	55
	Example 5.	
	10,000 capsules, each containing 20 mg. of $2-[\alpha-(3'-benzoylphenyl)]$ propionamide]-4-methyl pyridine hydrochloride were prepared as follows:	
	. 23	

	1,436,502	4
	Formulation:	
	2-[α-(3'-benzoylphenyl)propionamide]-4-methyl pyridine hydrochloride	
	Lactore 200 g.	
5	Com Starch 650 g.	
	Colloidal silicon oxide 485 g.	5
	Preparation:	
	The 2-[o-(3'-henzovinho1)	
_	The 2- $[\alpha$ -(3'-benzoylphenyl)propionamide]-4-methyl pyridine hydrochloride was mixed with the lactose, corn starch and colloidal silicon oxide and the resulting mixture was filled into 10,000 hard gelatin capsules of an appropriate α -constant.	
10	ture was filled into 10,000 hard gelatin capsules of an appropriate size.	
		10
	Example 6.	٠
	10,000 tablets, each containing 20 mg. of 2-[\alpha-(3'-benzoylphenyl)propionamide]- thiazole, were prepared as follows:	
	*	
	Formulation:	
15 -	$2-[\alpha-(3'-benzoylphenyl)$ propionamide]-thiazole 200 g.	15
	Lactose 1.590 g. Microcrystalline cellulose 600 g.	15
	Carboxy methyl storch	
	Colloidal silicon oxide	
20	Hydroxypropylcellulose g.	20
	Magnesium stearate 2 g.	20
	Preparation:	
•	The finely micronized 2-for/2/ honoral-board and	•
	The finely micronized 2-[\alpha-(3'-benzoylphenyl)propionamide]-thiazole and 2 g. of colloidal silicon oxide were granulated with the hydroxypropylcellulose dissolved in a 50:50 mixture of water and ethanol in a high provided him.	
25		
:	granulate was dried, passed through a 30 mesh screen and mixed with the lactose,	25
	16 mesh screen, mixed with the rest of the mean precompressed, passed through a	
30		20
		30
	Example 7.	•
:	1000 suppositories, each containing 25 mg. of $2-[\alpha-(3'-benzoylphenyl)]$ propionamide]-4-methyl pyridine hydrochloride, were prepared as follows:	
35	Formulation:	25
	2-[a-(3'-benzoyl phenyl)propionamide]-4-methyl	35
:	pyridine hydrochloride 250 g. Adep solidus 16.250	• • •
	16.250 g.	• •
	Preparation:	
10	The adeps solidus was melted in an electrically heated thermostatically controlled	40
		70
	amide]-4-methyl pyridine hydrochloride was added with constant stirring. The resulting suspension was poured into suppository moulds forming 1000 suppositories, each weighing approximately 1650 mg.	•
• •	weighing approximately 1650 mg.	. ~
15	WHAT WE CLAIM IS:—	45
	1. A compound corresponding to the general formula (I):	
:	R	•
	CO ZR ₁	
•	O = O = CON (I)	.*
	~ R ₂	
	where R represents a hydrogen atom lower ell-1 (C. C.)	
	where R represents a hydrogen atom, lower alkyl (C ₁ —C ₅) radical or cycloalkyl radical; R ₁ represents a hydrogen atom or lower (C ₁ —C ₃) alkyl radical; and R ₂ represents a heterocyclic group baying one or more heterocyclic group baying and the head of the heterocyclic group baying and the head of the hea	•
D	a heterocyclic group having one or more heteroatoms, or R_1 and R_2 represents	50
	The state of the s	50

	137705702	5
	the adjoining nitrogen atom form 3-oxo-4,5-benzo-1,2-thiazolinyl-1,1-dioxide, or a pharmaceutically acceptable salt thereof. 2. A compound as claimed in claim 1, wherein R represents a hydrogen atom or a methyl group.	
5	3. A compound as claimed in claim 1 or 2, wherein R ₁ represents a hydrogen atom. 4. A compound as claimed in any of claims 1 to 3, wherein R ₂ represents 2-thiazolinyl, 4-methylpyridyl, 3-hydroxypyridyl, pyridyl, 1,5-dimethyl-2-phenyl-pyrazolonyl or thiazolyl.	. 5
	5. A compound as claimed in claim 1, substantially as herein described with reference to Example 3	
10	to Dimitible 1.	
	6. A compound as claimed in claim 1, substantially as herein described with reference to Example 4.	10
15 .	7. A process for preparing a compound of formula (I) as defined in claim 1, which comprises hydrolysing a 3-benzoylphenyl α-substituted acetonitrile to form the corresponding alkanoic acid, converting the acid to an active derivative, and reacting the active derivative with an amine to form the desired amide derivative of formula (I). 8. A process as claimed in claim 7, wherein the active derivative is an acid chloride.	15
20	9. A process as claimed in claim 7, substantially as herein described with reference to the specific Examples 3 or 4	
20	10. A compound as claimed in claim 1, when produced by a process as claimed in any claims 7 to 9.	20
25	11. A pharmaceutical composition comprising a compound as claimed in claim 1 together with a non-toxic pharmaceutically acceptable carrier or diluent therefor. 12. A pharmaceutical composition as claimed in claim 11, substantially as herein described with reference to any of the preside line of the control of the contr	25
	described with reference to any of the specific Examples 5 to 7.	25

ELKINGTON AND FIFE, Chartered Patent Agents, High Holborn House, 52/54 High Holborn, London WC1V 6SH. Agents for the Applicants.

Printed for Her Majesty's Stationery Office by the Courier Press, Learnington Spa. 1976. Published by the Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.